Importance of Mangrove Carbon for Aquatic Food Webs in Wet–Dry Tropical Estuaries

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Received: 10 January 2014/Revised: 30 March 2014/Accepted: 3 April 2014/Published online: 23 April 2014 © Coastal and Estuarine Research Federation 2014

Abstract Mangroves are traditionally considered to provide important nutrition to tropical estuarine consumers. However, there is still controversy about this, and the extent and importance of these inputs are largely unquantified. In particular, there is no information for food webs of small estuaries that dominate wet-dry tropical coasts, where freshwater inflow is intermittent, leading to highly seasonal inputs of nutrients from terrestrial systems. Since the relative importance of the different sources depends on the type and extent of different habitats and on hydrological and topographic conditions, results from other regions/type of systems cannot be extrapolated to these estuaries. Here, δ^{13} C is used to determine the importance of mangrove-derived carbon for Penaeus merguiensis (detritivore; shrimp), Ambassis vachellii (planktivore; fish), and Leiognathus equulus (benthivore; fish) from six small wet-dry tropical estuaries that differ in mangrove (C_3) cover and in type of terrestrial vegetation adjacent to the estuary. Bayesian mixing models confirmed that mangrove material was important to consumers in all estuaries. There was a gradient in this importance that agreed with the extent of mangrove forests in the estuaries, as C3 sources were the most

Electronic supplementary material The online version of this article (doi:10.1007/s12237-014-9817-2) contains supplementary material, which is available to authorized users.

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important contributors to animals from the three estuaries with the greatest (>40 %) mangrove cover. There was also evidence of incorporation of C_3 material for the three estuaries with lower (<30 %) mangrove cover. Since these latter estuaries had no adjacent terrestrial C_3 forests, the detected C_3 influence can only be of mangrove origin. This shows that mangroves are important contributors to these food webs, underlining the importance of mangroves in supporting estuarine nursery ground value and fisheries productivity.

Keywords Bayesian mixing models $\cdot \delta^{13}C \cdot Estuaries \cdot Food$ webs \cdot Mangroves \cdot Stable isotope analysis

Introduction

The availability of adequate food and nutrients is vital to sustain the natural dynamics of biotic communities. In estuarine systems, animal communities generally rely on a combination of aquatic (autochthonous) and terrestrial (allochthonous) sources, with the importance of each source depending on the relative availability of material from the different origins (Polis et al. 1997; Bouillon et al. 2004; Abrantes et al. 2013). Given the present trend of increasing modification of estuaries and their catchments throughout the world (Lotze et al. 2006; Sheaves et al. 2014), it is crucial to understand the relative importance of these contrasting sources for estuarine consumers. However, the balance of contributions from different sources is still poorly resolved for many systems, especially in the tropics; while some studies suggest that terrestrial and mangrove/salt marsh material can be important (e.g., Wai et al. 2011; Abrantes et al. 2013), others have failed to find evidence of incorporation of mangrove/salt marsh or adjacent terrestrial vegetation and suggest that estuarine food webs are based on more easily

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assimilated aquatic producers (e.g., Chanton and Lewis 2002; Lin et al. 2007).

Although some research has been done for estuaries of large tropical rivers (e.g., Chanton and Lewis 2002; Abrantes et al. 2013), small perennial rivers in the wet tropics (e.g., Chong et al. 2001; Nyunja et al. 2009), near-pristine mangrove areas (Abrantes and Sheaves 2009a) and semiisolated estuarine floodplain pools (e.g., Abrantes and Sheaves 2008, 2010), information on the main sources of nutrition supporting consumers in the small estuaries that dominate wet-dry tropical coasts is still lacking. Given the widespread distribution of these systems, this presents a serious knowledge gap. The wet-dry climate is characterized by distinct wet and dry seasons, with most annual rainfall occurring during the wet season and very little falling during the dry season. Wet-dry tropical climate covers the coasts of most of northern Australia (from Central Queensland to northern Western Australia), eastern India, parts of Indo-China, eastern (Kenya to Mozambique and Madagascar) and western Africa (Senegal to Angola), Central (mostly in the western coast), Southern (parts of Brazil, Venezuela, and Colombia), and North America (parts of southern Mexico and Caribbean Islands) (Peel et al. 2007). Because there is considerable variation in assemblage structure (Sheaves and Johnston 2009) and because trophic processes in the different regions/ types of systems are likely to differ greatly depending on factors such as size, connectivity regimes, area of drainage basins, productivity of terrestrial and aquatic environments, and type and extent of adjacent vegetation (Abrantes et al. 2013), results from one region or one type of system cannot be extrapolated to other regions/systems without local validation.

Small estuaries are ubiquitous in the Australia's wet–dry tropics (Erskine et al. 2005; Sheaves et al. 2010), providing important feeding, spawning, and nursery habitats for a range of fish and invertebrates, many of recreational and/or economic importance (Beck et al. 2001; Dahlgren et al. 2006). Despite their small size and small catchments, these estuaries contribute disproportionately to coastal ecosystem functioning due to their prevalence and because they are generally less impacted by human activities than larger systems. Nevertheless, they are often subjected to specific local-scale impacts such as land-fill to convert mangrove areas to agricultural land and construction of barriers that restrict tidal/freshwater flows and limit connectivity of organic matter and animals between habitats (e.g., bund walls, levees, roads) (Boys et al. 2012).

Stable isotope analysis of carbon (δ^{13} C) is commonly used to study the importance of terrestrial vs. aquatic sources for estuarine food webs (e.g., Peterson et al. 1986; Bouillon et al. 2011). This is because different primary producers can have different δ^{13} C ratios (France 1996) and because δ^{13} C undergoes a small and predictable change from food source to consumer (0–1 ‰; DeNiro and Epstein 1978; McCutchan et al. 2003). However, estuarine producers can be spatially and/or temporarily variable in δ^{13} C values (Cloern et al. 2002; Guest et al. 2004); δ^{13} C of dissolved inorganic carbon (DIC) can affect the δ^{13} C of aquatic producers (Bouillon et al. 2008), and it can be methodologically difficult to appropriately sample aquatic producers such as phytoplankton and microphytobenthos (MPB), especially in turbid environments, meaning care is needed in the interpretation of stable isotope results.

We investigated δ^{13} C contributions to consumers in six estuaries in the wet-dry tropics of northern Australia to determine the relative importance of mangrove and terrestrial producers to the productivity of small wet-dry tropics estuaries. Systems surrounded by different combinations of mangrove forests and terrestrial vegetation with different mixes of C_3/C_4 plants provided an ideal situation to study the contribution of mangroves, because C₃ and C₄ sources are well separated in δ^{13} C (~-27 vs. ~-12 ‰) (Cerling et al. 1997). However, it can be difficult to differentiate contributions of organic matter imported from forests in the catchments from that of adjacent mangrove forests based on δ^{13} C alone, as both types of vegetation possess the same photosynthetic pathway (C_3) and are therefore characterized by similar δ^{13} C values. However, this complexity was used to advantage in the present study, as the systems considered have very small catchments and are subjected to short (2-3 months), well-defined wet seasons separated by long dry seasons, when freshwater flow is mostly absent (Finlayson and McMahon 1988). Consequently, they can receive terrestrial organic matter from the catchment for only 2-3 months of the year while for most of the year there is minimal potential for input of this material. Any incorporation of C₃ material detected at the end of the dry season is therefore likely to be of mangrove origin, rather than from adjacent terrestrial forests. Comparisons of contributions of C₃ sources between dry and wet seasons thus give further information on the input of mangrove vs. terrestrial material from the catchments.

Methods

Study Sites

The estuaries of six creeks spanning ~120 km of North Queensland's wet–dry tropics (Australia) were sampled: Sandfly, Cocoa, Doughboy, Crab, Mud, and Hell Hole Creeks (Fig. 1). These creeks are typical of the region: they are relatively short (<10 km navigable length) and narrow, with maximum mouth widths between 25 and 60 m, narrowing down to 5–15 m at the upper limits of navigation. All systems are tide dominated, with tidal incursions ranging between 2 and 5 km. Tides are semi-diurnal with a maximum range of ~4 m. Depths at low tide are typically ~3.5 m closer to the mouth, decreasing with distance upstream until the



Fig. 1 Map showing the study sites in North Queensland, Australia

limits of navigation. The substrates are dominated by sand and mud. Macroalgae are very rare and seagrass only occurs in the mouths of Cocoa and Crab Creeks. The climate of the area is characterised by a short rainy season from December to March and a long dry season from April to November (Fig. 2), when there is little or no freshwater inflow from intermittent feeder streams, leading to strong seasonality in potential inputs of nutrients from terrestrial systems.

Mangrove forests occur at the lower reaches of all estuaries, with a mangrove fringe at least 5 m wide also present through most of their length. Salt marshes (dominated by the C_4 grass *Sporobolus virginicus*) and saltpans sometimes occur adjacent to the mangrove forests, in the upper intertidal. These areas are generally flooded only on the highest spring tides and for relatively short periods. In Doughboy, Mud, and Hell Hole Creeks, surrounding vegetation is mostly mangrove and terrestrial forests, whereas at Crab and Sandfly Creeks, the adjacent area is dominated by C₄ producers (Table 1), mostly salt marsh plants but also including pasture grass and, in Sandfly Ck, limited horticulture. In Cocoa Ck, the surrounding area is dominated by salt flats (Table 1). Cocoa and Sandfly Creeks have U-shaped cross-channel profiles with steep banks that provide little area for benthic microalgae production; Crab Ck has a slightly larger intertidal area, followed by Doughboy Ck, which has areas of intertidal banks (up to ~ 3 m wide) suitable for benthic productivity. Hell Hole and Mud Creeks are shallower, with large intertidal areas, and Mud Ck has the largest intertidal area of all estuaries (Table 1).

Sample Collection and Analysis

Juveniles of two fish and one penaeid prawn species were sampled at each estuary in mid-March 2008, just after the 2007/2008 wet season, and again in mid-November 2008, at the end of the dry season (to represent carbon accumulated during the wet and dry seasons, respectively). The only exception was Sandfly Ck, which could not be sampled in the wet season due to local flooding. Species considered were the Vachell's glassfish *Ambassis vachellii*, the common ponyfish *Leiognathus equulus*, and the banana prawn *Penaeus merguiensis*. *A. vachellii* is an estuarine spawner, and individuals captured were likely to have spent their whole life in the estuary of capture; *L. equulus* and *P. merguiensis* are offshore spawners, with juveniles (*L. equulus*) or postlarvae (*P. merguiensis*) recruiting into estuaries at small sizes





Table 1 Turbidity, salinity, intertidal area relative of total estuary surface area (ranked from 1 to 5), and percentage cover of mangroves, terrestrial C_3 forests, total C_3 cover (including mangroves and terrestrial forests), C_4 vegetation (including salt marsh, pasture land, and sugarcane plantations), savannah vegetation (C_3/C_4 mix), and salt flats in each estuary.

Percentage cover calculated for the area within 1 km from the river margins, up to the upper extend of tidal reach. For turbidity and salinity, data for the wet and dry seasons are presented, and values indicate measurements made at the lower (close to estuary mouth) followed by the upper reaches (close to the limit of saltwater intrusion)

Estuary	Salinity		Turbidity (NTU)		Intertidal	% Cover						
	Wet	Dry	Wet	Dry	(rank)	Mangrove (C ₃)	Terr C ₃ forest	Tot C ₃	C ₄	Savanna	Salt flat	
Cocoa Ck	34/35	39/47	78/37	20/17	1	18.9	0.0	18.9	5.9	20.5	54.7	
Crab Ck	33/32	40/43	121/28	12/18	2	27.6	0.0	27.6	36.6	0.6	35.2	
Doughboy Ck	26/30	46/45	72/43	15/21	3	49.6	32.0	81.6	15.6	2.3	<1	
Hell Hole Ck	16/18	42/37	25/18	8/28	4	41.5	14.1	55.5	<1	43.9	<1	
Mud Ck	28/32	43/46	26/42	12/19	5	47.4	20.6	68.0	19.9	4.5	7.6	
Sandfly Ck	-	40/45	_	30/56	1	20.0	0.0	20.0	43.1	24.8	12.0	

(<20 mm for *L. equulus* (Sheaves et al. 2013); <3-mm carapace length for *P. merguiensis* (Haywood and Staples 1993)) early in the pre-wet season, so animals collected would likely have spent a minimum of a few months in the systems before both sampling occasions (Robertson and Duke 1990; Haywood and Staples 1993).

P. merguiensis juveniles are mostly detritivores, feeding on large amounts (up to 75 %) of flocculent detritus (including mangrove material) and also on small invertebrates such as crustaceans (ostracods, calanoid copepods, brachyuran larvae) and gastropods (Robertson 1988). *A. vachellii* feeds mostly on planktonic crustaceans such as decapoda larva, copepods, and amphipods (Wilson and Sheaves 2001). *L. equulus* juveniles have a more benthic-associated diet, feeding mostly on small benthic prey (harpacticoid copepods, gammarid amphipods, gastropods, bivalves, polychaetes) and to a lesser extent on pelagic prey such calanoid copepods (Wilson and Sheaves 2001; Mavuti et al. 2007).

These three species were selected because they are abundant and ubiquitous in Australia's wet-dry tropics estuaries (Sheaves and Johnston 2010; Sheaves et al. 2012), represent contrasting trophic pathways and because they are fast growing (Robertson and Duke 1990; Haywood and Staples 1993), meaning that their stable isotope composition should accurately reflect the diet over the last 1-2 months. After a change in diet, it takes some time for the isotopic composition of consumers to reflect the new diet, depending both on growth and on metabolism. For muscle of small (<5 g), growing fish and invertebrates such as those used in the present study, carbon half-lives are less than 1 month (Guelinckx et al. 2007; Weidel et al. 2011). Thus, the stable isotope composition of these animals provides time-integrated information on the most important sources of nutrition for food webs at each estuary, and the ~3.5-month time lag between the beginning of the 2007/2008 wet season and the March 2008 sampling, and the ~8-month time lag between the end of the 2007/2008 wet season and the November 2008 sampling (see Fig. 2) means that the δ^{13} C values of animals at the times of capture will reflect their diet in wet and dry conditions, respectively.

Animals were captured with a 5-mm mesh monofilament drawstring cast net deployed from a small boat. Sampling was done over the low tide period, when mangrove forests were drained and animals forced into the channels. Each estuary was sampled at its lower (close to the mouth), middle, and upper reach (close to the limit of saltwater intrusion). Whenever possible, 15 individuals of each species (of similar sizes) were collected from each reach. Samples were stored frozen until processing. Salinity and turbidity were also measured on each sampling occasion. Salinity was measured using an optical salinometer (accuracy ~1) and turbidity with a TPS WP-88 handheld turbidity meter (accuracy ~1 NTU). Instruments were referenced to standards before each sampling trip. In the laboratory, animals were identified and measured (standard length for fish; total length for prawns), and white muscle tissue was excised from the trunk below the dorsal fin of fish and from the abdominal muscle of prawns. Samples were then dried for 48 h at 60 °C and homogenized into a fine powder with a mortar and pestle. For each species, similar amounts (by weight) of dried tissue from each of the individuals collected at each reach/ estuary were combined into a single sample to reduce the effect of intraspecific variability providing the best possible estimate of carbon isotopic composition of a species in each sample (Lancaster and Waldron 2001). This material was then homogenised by manually shaking the vials, and ~ 0.1 g was subsequently weighed into 8×5 tin capsules. The carbon stable isotope composition of each sample was measured with an Isoprime isotope ratio mass spectrometer (IRMS) coupled with an element analyzer. Results are expressed as per mil (‰) deviations from the standards, as defined by the equation: $\delta^{13}C{=}[(\delta^{13}C_{\text{ sample}}\,/\,\delta^{13}C_{\text{reference}}){-}\,1]{\times}10^3$ and had a precision of ± 0.1 ‰ (SD), calculated from multiple runs of the same prawn and flour samples. No corrections for lipid content were made as C-N ratios of all samples were <3.5 (Post et al. 2007).

Data Analysis

For each estuary, Bayesian mixing models were used to quantify the contribution of the main classes of producers to each species, using SIAR (Stable Isotope Analysis in R; Parnell et al. 2010). Because the stable isotope composition of animals from the three reaches within an estuary often differed, models were run for each reach separately. Since generally only one sample (composed of up to 15 individuals, pooled) was available for each reach, the siarsolo command was used (Parnell et al. 2010). In some cases, however, two or three composite samples were available for the same reach and season, in which cases δ^{13} C results were averaged (arithmetic mean) between samples and the mean used in the model. Bayesian mixing models produce a range of solutions given the available sources while taking into account uncertainty and variation in consumer stable isotope composition and trophic enrichment factors (TEF). A δ^{13} C TEF of 1.0 ‰ was used, while taking into account the different species' trophic level, as appropriate for non acid-treated muscle tissue (McCutchan et al. 2003), and a TEF standard deviation (SD) of 1.5 ‰ was used to account for the uncertainty in this value (Vander Zanden and Rasmussen 2001; McCutchan et al. 2003; Caut et al. 2009). P. merguiensis juveniles were considered to be of trophic level 2.5 (Robertson 1988; Abrantes and Sheaves 2009b), and A. vachellii and L. equulus juveniles of trophic level 3 (Wilson and Sheaves 2001; Mavuti et al. 2007). Concentration dependencies were set to zero. Because SIAR is sensitive to variation in discrimination factors (Bond and Diamond 2011), a sensitivity analysis was done in which additional models were run using TEFs of 0 and 2 ‰, to determine if different scenarios would lead to different final results regarding contribution of C₃ sources.

Potential sources considered were C₃ producers (which include mangroves), C₄ producers (which include salt marsh), plankton, and microphytobenthos (MPB). For Cocoa and Crab Creeks, seagrass was also considered as a potential source, as seagrass beds occur in the mouths of these estuaries. Primary producers were not collected, so the δ^{13} C values used in the models were based on data from similar systems of the region or data from the literature. For plankton, the δ^{13} C value used was -20.5 ‰, and for MPB, a value of -14.0‰ was used, based on the average δ^{13} C of plankton/MPB collected from 15 small estuaries spanning over 600 km of the wet-dry tropical Queensland coast (own unpublished data). For seagrass, the average value of -11.5 ‰ was considered, based on the review by Hemminga and Mateo (1996) on the variability in seagrass δ^{13} C. For C₃ sources, a δ^{13} C of -27 ‰ was used, and for C_4 sources -12 ‰ was used (Cerling et al. 1997). Because these values were taken from other studies, large source SDs of 2 ‰ were used to account for the uncertainty. For MPB, a larger SD of ± 3 ‰ was used, as MPB can have a relatively large variability in δ^{13} C in North Queensland estuaries (own unpublished data). These large source SDs, coupled with the large TEF SDs used (1.5 ‰), should lead to conservative results regarding the importance of the different sources. Note that even if source values are not precise, all models were calculated based on the same values so results will be comparable among the sites. Because only one element was used and the number of sources was 4–5, we did not expect to be able to distinguish between the contributions of all sources. However, our aim was to identify and quantify the importance of C₃ mangroves, and since C₃ producers are well separated from the remaining sources, their contribution can be estimated with confidence, especially since the three consumer species often had δ^{13} C that could only be explained by some contribution of C₃ sources.

The proportion of mangrove forests, terrestrial forests, C₄-dominated vegetation (including salt marsh and cattle pastures), saltpan, and savanna vegetation (mix of C₃ and C₄ vegetation) in the area adjacent to each estuary were estimated using SigmaScan Pro, based on freely available images from Google Earth and supported by detailed ground truthing. Because the catchments of these systems have not been delineated, the percentage of vegetation types within 1 km of estuary margins was considered as indicative of adjacent available producers. To help separate contributions of carbon from mangrove origin from that of terrestrial C₃ forests, the effects of the relative mangroves cover and of overall C3 cover (includes both mangroves and C₃ terrestrial forests) on the contribution of C3 sources to consumers were modeled for each species and season using multiple regression models with backward elimination. The aim was to determine the extent to which contribution of C3 sources (%; based on Bayesian mixing models) (dependent variable) can be explained by mangrove cover alone, and whether total C₃ cover (i.e., including also terrestrial forests) provides more explanatory power.

Classification and regression tree analyses (CARTs; De'ath and Fabricius 2000) were used to explain the extent to which the importance of C₃ sources depends on estuary, reach, species, and season. Input data were the modal contributions of C₃ sources for each group (Parnell et al. 2010). CART analysis is robust non-parametric test that successively splits the dataset into two relatively homogeneous and mutually exclusive groups based on minimising the within-group sum of square residual deviation. The trees are represented in a graphical way, with the root node on top, representing the initial assemblage of data, from which the branches and leaves emerge. Splits close to the node are more important than those at the bottom of the tree, providing greater improvement to the fit of the model. The relative lengths of the vertical lines associated with each split gives indication of the proportion of the total sum of squares explained by each split. The size of the tree (or number of leaves), corresponding to the final number of groups, was selected by tenfold cross-validation

and the 1-SE tree, i.e., the smallest tree with cross validation error within 1 SE of the tree with the minimum cross validation error, was selected as the final tree model (De'ath and Fabricius 2000). Analyses were conducted using the Trees Plus package (De'ath and Fabricius 2000). Because not all species occurred at all reaches and at all seasons and this could hinder the identification of seasonal effects by the CART, seasonal differences in importance of C3 sources were further investigated for pairs of species that occurred in the same estuary and same reach for both seasons. Although wet season samples were collected before the dry samples, in March and November 2008, respectively, results are presented as changes in C₃ contribution from the dry to the wet season, to facilitate interpretation of the effect of the wet season. The presence of seasonal shifts in importance of C3 sources was tested using CART analysis where the dependent variable was the difference in mode of contribution between seasons, and the independent variables were species, estuary, reach, and season. The input data consisted of zeroes for the dry season, i.e., the starting point against which the effect of the wet season was measured, and input values for the wet season corresponded to the differences in mode of contribution between the two seasons. A split between seasons with zero in the dry and the difference in mode contribution in the wet season would indicate a significant seasonal change in importance of C₃ sources, while the lack of a split would indicate that the importance C₃ sources was similar for the two seasons.

Results

Environmental Parameters

For all estuaries, salinities were lower during the wet season than in the dry season (Table 1). In the wet season, salinities were generally similar and close to sea water in with the exception of Hell Hole, where waters were less saline (16-18). Within each estuary, salinities did not vary by much between the lower and upper reaches (maximum difference between reaches only 4; Table 1). In the dry season, however, salinities were higher than seawater for all estuaries (range, 37–47). The greatest difference in salinity between upper and lower reaches occurred in Cocoa Ck (difference of 8), while for the remaining estuaries, the upstream-downstream differences were <5. In general, salinities were higher at the upper reaches than at the lower reaches, with the exception of Doughboy Ck (similar salinity at both reaches) and Hell Hole Ck (salinity at lower reach higher than at upper reach). There were often differences in turbidity between the lower and upper reaches (Table 1). Wet season turbidities varied between lower reaches of estuaries, with clearer waters in Hell Hole and Mud Creeks (25 and 26 NTU, respectively),

intermediate turbidity in Cocoa and Doughboy Creeks (78 and 72 NTU, respectively), and a maximum of 121 NTU at Crab Ck. In the upper reaches, turbidity levels were low and more similar between estuaries, ranging from 18 to 43 NTU. In the dry season, turbidities were similar among estuaries and generally low, between 8 and 30 NTU in the lower reaches and between 17 and 56 NTU in the upper reaches (Table 1).

Animal δ^{13} C and Mixing Model Results

For all three species, there were differences in δ^{13} C between estuaries and between seasons (Table 2; Fig. 3). In general, animals from Mud, Doughboy, and Hell Hole Creeks had the lowest δ^{13} C values and those from Cocoa, Crab, and Sandfly Creeks the highest (Fig. 3). In the wet season, all three species had relatively similar δ^{13} C values within each estuary, but in the dry season, the three species often differed in δ^{13} C (Fig. 3). Moreover, while the δ^{13} C values of a species were similar for the three reaches during the wet season, in the dry season, those values generally differed between reaches, often by more than 3 ‰ (Fig. 3). Accordingly, mixing model results show that, within each estuary, the three species depended on a similar combination of sources in the wet season, while in the dry season, the three species reliance on the different sources varied (Fig. 4; Electronic Supplements 1 and 2). Additionally, for each species, the contribution of the different sources was similar between reaches for the wet season but generally differed between reaches in the dry season (Fig. 4; Electronic Supplements 1 and 2).

Mixing models based on different TEFs (0, 1, and 2 ‰) lead to similar patterns of spatial and temporal variability in importance of C₃ sources for the three species. Overall, C₃ material was an important source for the three species in all estuaries, but this importance varied between estuaries, reaches, seasons, and species (Electronic Supplement 1 and 2). For example, when considering a TEF of 1 ‰, C₃ contributions varied from 5 to 41 % (95 % credibility interval (CI)) for P. merguiensis from the downstream reaches of Cocoa Ck, to 70-97 % for L. equulus at the upstream reach of Mud Ck (Fig. 4; Electronic Supplements 1 and 2). Models run using TEFs of 0 and 2 ‰ led to similar results: C₃ contribution was the lowest for P. merguiensis from the downstream reaches of Cocoa Ck (95 % CI=1-24 ‰ and 10-48 ‰ when considering TEFs of 0 and 2 ‰, respectively), and the highest for L. equulus at the upstream reach of Mud Ck (95 % CI=58-95 ‰ and 76–98 ‰) (Electronic Supplement 2).

In general, C₃ sources were the most important contributors for animals in the three estuaries with the highest mangrove cover (Doughboy, Hell Hole, and Mud Creeks), while in estuaries with lower mangrove cover (Cocoa, Crab, and Sandfly Creeks), animals relied on a more balanced combination of terrestrial and aquatic sources, including benthic and

Table 2 Size range and carbon stable isotope composition (mean \pm SD) of each species in the wet (March 2008) and dry seasons (November 2008)

	Wet Season			Dry Season			
	Size (mm)	$\delta^{13}C$	п	Size (mm)	$\delta^{13}C$	п	
P. merguiensis							
Cocoa Ck	35–45	-18.3 ± 1.0	L:1(15); M:1(15); U:1(15)	30–50	-17.7 ± 1.1	L:1(15); M:1(3)U:1(15)	
Crab Ck	35–45	-20.0 ± 1.1	L:1(16); M:1(15); U:1(14)	35–45	-18.3 ± 0.7	L:1(13); M:1(3); U:1(4)	
Doughboy Ck	35–45	-23.0 ± 0.6	L:1(15); M:1(15); U:1(15)	30–50	-21.6 ± 0.5	L:NC; M:2(4-8); U:1(5)	
Hell Hole	35–45	-22.4 ± 0.1	L:1(10); M:NC; U:1(7)	35–45	-21.6 ± 3.1	L:1(5); M:1; 7); U:1(15)	
Mud Ck	35–45	-23.7 ± 0.3	L:1(15); M:1(15); U:1(15)	25-40	-22.2 ± 1.9	L:1(5); M:1(13); U:1(7)	
Sandfly Ck	_	_	-	30–50	-18.2 ± 0.7	L:1(15); M:1(15); U:1(13)	
A. vachellii							
Cocoa Ck	35–45	-18.2 ± 0.4	L:NC; M:2(5-15); U:NC	35-60	$-18.0 {\pm} 0.8$	L:1(7); M:NC; U:1(2)	
Crab Ck	35–45	-20.0 ± 0.3	L:1(15); M:1(15); U:1(15)	40–45	-21.4 ± 1.9	L:1(1); M:1(1)	
Doughboy Ck	35–45	-21.8 ± 0.5	L:1(15); M:1(15); U:1(15)	_	_	_	
Hell Hole	35–45	-21.9 ± 0.4	L:1(15); M:1(15); U:1(15)	40–50	-21.7 ± 0.4	L:NC; M:3(3-8); U:NC	
Mud Ck	35–45	-22.5 ± 0.6	L:1(15); M:1(15); U:1(15)	35-50	-24.4 ± 0.3	L:3(5); M:NC; U:NC	
Sandfly Ck	_	-	_	35–40	-21.3 ± 0.5	L:1(15); M:1(15); U:1(9)	
L. equulus							
Cocoa Ck	30–45	-17.7 ± 0.8	L:3(6-7); M:1(7)	55-75	-20.6 ± 3.5	L:1(4); M:1(1); U:NC	
Crab Ck	30–40	-19.8 ± 0.3	L:NC; M:1(2); U:1(8)	35-50	-21.8 ± 1.0	L:NC; M:2(4-5); U:1(5)	
Doughboy Ck	30–40	-23.0 ± 0.6	L:1(7); M:1(15); U:1(15)	35–50	-24.7	L:NC; M:NC; U:1(1)	
Hell Hole	25-40	-22.5 ± 0.5	L:1(5); M:1(7); U:1(7)	45–55	-23.3 ± 2.0	L:2(2-3); M:1(3); U:NC	
Mud Ck	30–45	-23.5 ± 1.5	L:1(14); M:1(7)U:1(11)	45-70	-25.9 ± 2.6	L:1(8); M:1(7); U:1(15)	
Sandfly Ck	_	_	_	15-50	-19.7 ± 1.2	L:1(10); M:1(13); U:1(15)	

n is number of samples for the lower (L), middle (M), and upper (U) reaches, followed (in parenthesis) by the number of individuals pooled in each sample. NC not collected

planktonic algae (Electronic Supplements 1 and 2). In these latter estuaries, when considering a TEF of 1 ‰, C₃ sources contributed to all species in all reaches, with lower bounds of the 95 % CI \geq 10 % in 29 out of the 37 cases, >20 % in 9, and >30 % in 2 cases (Electronic Supplements 1 and 2). When

considering a TEF of 0 ‰, the lower bounds of the 95 % CI were >10 % for 14 out of the 37 cases (>20 % in two cases), and for models run using TEFs of 2 ‰, lower bounds of the 95 % CI were >10 % in 26 out of the 27 cases (>20 % in 22 cases and >30 % in 5 cases) (Electronic Supplement 2).

Fig. 3 Carbon stable isotope composition of consumers. The δ^{13} C values of possible sources (mean ± SD; as used in the Bayesian mixing models) are also indicated below the plots (see text for details). C3=C₃ sources; C4=C₄ sources, MPB = microphytobenthos; Plk = plankton; SG = seagrass (only present in Cocoa and Crab Creeks)



Fig. 4 Relationships between the estimated mangrove cover in the vicinity of each estuary and the mode contribution of C3 sources (based on Bayesian mixing models, while considering a TEF of 1 ‰) to each species. Sites are represented by the estimated mangrove cover: Cocoa Ck, 18.9 %; Sandfly Ck, 20.0 %; Crab Ck, 27.6 %; Hell Hole Ck, 41.5 %; Mud Ck, 47.4 %; Doughboy Ck, 49.6 %. All relationships were significant (p < 0.05). See Table 3 for relationship details



For both seasons, there were positive relationships between the relative area of mangrove cover (in %) and the modal contribution of C_3 sources for the three species (Table 3; Fig. 4). These relationships were present when models were run using TEFs of 0, 1, and 2 ‰ (Table 3). Backward multiple linear regressions show that mangrove cover was the most important factor explaining the importance of C_3 sources to consumers, and that including terrestrial C_3 forest cover (to make total C_3 cover) in the models did not improve explanatory power in any case (Table 3). The only exception was for *A. vachellii* in the dry season, for which no effect of mangrove or total C_3 cover was detected for models run using a TEF of 0 ‰, as all variables were removed from the regression equation (Table 3).

For the CART analyses, models run using TEFs of 0, 1, and 2 ‰ led to similar results (Fig. 5). In the three cases, four-leaf CARTs, explaining 63, 59, and 65 % of the total variability, respectively, indicate that the contribution of C_3 sources is primarily dependent on estuary, as material of C_3 origin was more important for consumers in Doughboy, Hell Hole, and Mud Creeks, the estuaries with higher (>40 %) mangrove cover, than in Cocoa, Sandfly, and Mud Creeks, the estuaries with lower (<30 %) mangrove cover (Fig. 5). This first split in the data explained most of the total variability: for the model run using a TEF of 0 ‰, it explained 46 % of the variability,

while for the models run on TEF of 1 and 2 ‰, it explained 45 and 50 %, respectively. Although there were small differences between models in the lower branches of the trees, the three models agree that for the three creeks with highest mangrove cover (Doughboy, Hell Hole, and Mud Creeks), the contribution of C_3 sources was greater for *L. equulus* than for *P. merguiensis* and *A. vachellii* (Fig. 5).

There was also evidence of seasonal differences in importance of C₃ sources, although this varied between species (Fig. 6). CARTs based on solutions of the mixing models run using different TEFs led to similar results. When a TEF of 0 % was used in the mixing models, the resulting three-leaf CART indicates a significant effect of season but only for L. equulus, for which the importance of C_3 sources was greater in the dry season than in the wet season (Fig. 6a). CARTs based on mixing models with higher TEFs of 1 and 2 ‰ also showed a similar effect for A. vachellii, while for P. merguiensis, C₃ sources were generally more important in the wet season, especially for the middle and lower reaches of estuaries (Fig. 6a, b) where the largest mangrove areas were generally present and regularly submerged. Therefore, while for the two fish species, the importance of C₃ sources was greater in the dry season, for the prawn species C₃ sources were more important in the wet season.

Table 3 Results from stepwise multiple linear regression analysis testing the effects of mangrove cover and total C₃ cover (in %) on the modal contribution of C₃ sources (C_{3-cont}; based on Bayesian mixing models) for *Penaeus merguiensis*, *Ambassis vachellii*, and *Leiognathus equulus* in the wet and dry seasons, while considering TEFs of 0, 1, and 2 %. R^2 and *p* values are presented for the variables included in the models

		Mangro	ve	Total	C ₃		
		R^2	p level	R^2	p level	Equation	
P. merguiensis							
Wet season	TEF=0‰	0.872	< 0.0001	_	_	$C_{3-cont}=0.65 \times mang cover+12.83$	
	TEF=1‰	0.808	< 0.0001	_	_	C _{3-cont} =0.83×mang cover+14.11	
	TEF=2‰	0.841	< 0.0001	_	_	C _{3-cont} =1.08×mang cover+16.09	
Dry season	TEF=0‰	0.520	0.0011	_	_	C _{3-cont} =0.53×mang cover+13.74	
	TEF=1‰	0.448	0.0033	_	_	C _{3-cont} =0.69×mang cover+14.98	
	TEF=2‰	0.472	0.0023	-	-	C _{3-cont} =0.78×mang cover+18.17	
A. vachellii							
Wet season	TEF=0‰	0.892	< 0.0001	_	_	C _{3-cont} =0.43×mang cover+18.11	
	TEF=1‰	0.788	< 0.0001	_	-	C _{3-cont} =0.67×mang cover+18.89	
	TEF=2‰	0.812	< 0.0001	_	_	C _{3-cont} =0.84×mang cover+27.24	
Dry season	TEF=0‰	_	_	_	_	-	
	TEF=1‰	0.528	0.0172	_	-	C _{3-cont} =0.93×mang cover+19.60	
	TEF=2‰	0.484	0.0376			C _{3-cont} =0.92×mang cover+33.05	
L. equulus							
Wet season	TEF=0‰	0.794	0.0001	_	-	C _{3-cont} =0.71×mang cover+10.55	
	TEF=1‰	0.680	0.0010	_	-	C _{3-cont} =0.97×mang cover+12.51	
	TEF=2‰	0.871	< 0.0001	_	_	C _{3-cont} =1.16×mang cover+17.65	
Dry season	TEF=0‰	0.667	0.0007	_	_	C _{3-cont} =0.97×mang cover+12.97	
	TEF=1‰	0.573	0.0044	_	_	C _{3-cont} =1.03×mang cover+22.01	
	TEF=2‰	0.762	< 0.0001	_	-	C _{3-cont} =1.11×mang cover+29.43	

Importance of Mangrove Carbon for Estuarine Food Webs

In general, results indicate that C₃ material is important for aquatic food webs in small wet-dry tropical estuaries. However, this importance differs between systems, depending on the type and extent of adjacent vegetation. C₃ contribution is correlated with the relative extent of mangrove forests at each estuary, and adding the terrestrial C₃ forest cover to the models did not provide greater explanatory power in any case, suggesting that mangroves, rather than terrestrial forests in the adjacent area, were the main sources of C₃ carbon for consumers in these systems. Indeed, even for Sandfly, Cocoa, and Crab Creeks, where terrestrial forests are absent, mangrove forests covered <30 % of the adjacent area, and where C₄ vegetation (including salt marsh, pasture land, and sugarcane plantations) and salt flats dominated the adjacent area, C₃ carbon was still important for consumers, with mode contributions always \geq 25 % and lower bounds of the 95 % CIs > 5 % in all but one case, when considering TEF 1 ‰ for example (see Electronic Supplement 1).

For all sites, C_3 sources had some importance for all species even in November 2008, 8 months after the end of the previous wet season, further indicating that even for the estuaries where adjacent C_3 forests are present, this C_3 input was from mangrove productivity rather than from forests in the adjacent catchment. Indeed, the minimal rainfall during the 2008 dry season (see Fig. 2) was unlikely to be sufficient to transport significant amounts of C3 terrestrial organic material into the waterways. The small catchments of these estuaries and little, if any, freshwater inflow during most of the year (Sheaves 1996) also limit the possibility that any substantial material from the upstream catchment is imported into the estuary. Although mangrove carbon is considered to be of poor nutritional quality, tropical mangrove forests are highly productive, and high quantities of nutrients, organic matter, and mangrove litter regularly enter these systems (Jennerjahn and Ittekkot 2002; Kristensen et al. 2008). Several invertebrate and fish species, including the species considered in the present study, move into mangrove forests at high tides for food and protection (Vance et al. 1996; Sheaves and Molony 2000). The relative importance of mangroves can be further increased in small estuaries such as those from the present study as these are narrow, with high ratios of mangrove area to open water area (Robertson and Blaber 1992). Thus, the often >50 % modal contribution of C3 sources in the three estuaries with higher mangrove cover, even in the dry season and even when a TEF of 0 ‰ was considered, indicates that mangroves can be the main sources of nutrients supporting food webs in these systems. If levels of aquatic productivity are similar for systems with and without extensive mangrove forests, it is likely











that estuaries with larger areas of mangrove forests can fuel more abundant consumer communities.

Sensitivity Analysis and Other Considerations

The use of different TEFs (0, 1, and 2‰) in the Bayesian mixing models led to similar patterns of spatial and temporal variation in importance of C₃ sources for the three consumer species. This sensitivity analysis confirms that there is an incorporation of C₃ material by these estuarine species and that there is seasonality in this importance, although results based on TEF of 2 ‰ led to stronger patterns than models based on TEF of 1 and 0 ‰ (i.e., greater importance of C₃ sources for all species and stronger seasonal effects). This was expected, given the low $\delta^{13}C$ of C₃ sources. Although the average value of δ^{13} C TEF found in the literature is generally <1 ‰ (e.g., Vander Zanden and Rasmussen 2001: 0.5±1.2 ‰ (±SD); Post 2002: 0.4±1.3 ‰ (±SD); McCutchan et al. 2003: 0.5 ± 0.13 ‰ (±SE); Caut et al. 2009: 0.8 ± 0.1 ‰ (±SE)), those values are based on meta-analyses that consider multiple taxa, environments, and tissues, and δ^{13} C TEF varies with all these factors (e.g., Vander Zanden and Rasmussen 2001; McCutchan et al. 2003; Caut et al. 2009). When considering only muscle tissue with no lipid removal, as in the present study, the average δ^{13} C TEF is higher: reviews by McCutchan et al. (2003) and Caut et al. (2009) showed that average δ^{13} C TEF for non-lipid treated muscle tissue was 1.1 ± 0.3 and 1.8 ± 0.8 ‰ (\pm SD), respectively. Other studies (not considered in those reviews) also found that a $\delta^{13}C$ TEF ≥ 2 ‰ is more appropriate for fish muscle (e.g., Barnes et al. 2007; Elsdon et al. 2010) and results from further studies indicate a δ^{13} C TEF of 2 ‰ or higher, despite that fish muscle did not reach equilibrium (Gorokhova and Hansson 1999; Guelinckx et al. 2007; Buchheister and Latour 2010). Similarly, for crustacean muscle, δ^{13} C TEFs larger than 0 ‰ have been reported (Yokoyama et al. 2005: 2.2 ‰; Suring and Wing 2009: 0.8 ‰). Therefore, results from the mixing models based TEF of 0 ‰ can be considered conservative regarding the contribution of C₃ sources, as higher TEF values lead to lower corrected δ^{13} C which in turn leads to higher contributions of C3 sources. Models based on TEFs of 1 and 2 ‰ can be considered closer to the reality in these systems.

It can be argued that the differences in importance of C_3 material between estuaries resulted from spatial differences in



Fig. 6 Classification and regression trees explaining the seasonal changes in importance of C_3 sources for the three consumer species when Bayesian models were computed using a TEF of a 0 ‰, b 1 ‰, and c 2 ‰. Explanatory variables were species, estuary, reach (lower, middle, and upper), and season. Models calculated based on the differences in mode contribution of C_3 sources (based on Bayesian mixing model results) between the dry and wet season using data from all cases when a species was collected in the same estuary and same reach at both seasons. Graphs below each leaf are the histograms of distribution of the values of shifts in contribution of C_3 sources (in %). Mean shift (in %) and sample size (in brackets) for each group are indicated below each graph

 δ^{13} C values of aquatic producers, which were not measured. Note however that estuarine aquatic producers are often temporarily and spatially variable in δ^{13} C at small scales (e.g., Cloern et al. 2002; Guest et al. 2004), so a sample collected at any point in time (or space) is unlikely to be representative of the source available throughout the area over time. This is especially the case for macrotidal systems such as the ones of this study. For example, in similar tropical small creeks, DIC- δ^{13} C varies up to ~10 ‰ with tidal level (Bouillon et al. 2007; Maher et al. 2013), and this would lead to similar changes in phytoplankton δ^{13} C in less than a day. However, because the six systems considered have similar conditions in terms of size, depth, tidal ranges, turbidity, climate, and hydrology, the average carbon stable isotope composition of the different aquatic primary producer categories (e.g., plankton, MPB) is likely to be similar between systems.

Although in the presence of mangroves, aquatic primary producers can have lower than expected $\delta^{13}C$ due to the incorporation of ¹³C-depleted DIC of mangrove origin (Bouillon et al. 2008; e.g., through flushing of crab burrows (Bouillon et al. 2007)); due to their small sizes, large tidal ranges (up to ~4-m semi-diurnal tides), and relatively shallow depths, the waters in these systems are likely to be well-mixed by tides, and the rapid water exchange is likely to minimize the effect of mangrove-derived ¹³C-depleted DIC over δ^{13} C of phytoplankton and other aquatic producers. For example, water residence time in a similar creek in southern Oueensland was of only ~1 tidal cycle despite a narrower tidal range (spring tides of ~2 m) (Maher et al. 2013), meaning it is likely that ¹³C-depleted DIC of mangrove origin is rapidly diluted and does not affect δ^{13} C of primary producers to the point of affecting δ^{13} C of secondary consumers. Although the relatively high salinities found in the dry season could be interpreted as resulting from low mixing of estuarine and marine waters, they are more likely to be a result of high evaporation rates over the mangroves, salt marshes, and salt flats (Ridd et al. 1997; Ridd and Stieglitz 2002). This phenomenon leads to short wet-dry tropical estuaries such as these rapidly becoming hypersaline over the whole length, even if there is effective tidal mixing (Ridd and Stieglitz 2002). Consequently, the time-averaged stable isotope composition of plankton and other primary producers such as MPB should be similar between systems.

A number of studies found strong variations in plankton/ seston δ^{13} C in estuaries, which were related to distance to mangroves (e.g., Hemminga and Mateo 1996; Bouillon et al. 2000). These studies were, however, done in large systems with high freshwater flows that discharge into large bays having, consequently, strong salinity gradients. It is well known that there is a strong positive relationship between salinity and $\delta^{13}C_{DIC}$ (e.g., Fry 2002; Gillikin et al. 2006), so it is likely that the distance to mangroves was not the only cause of those detected gradients in plankton/seston δ^{13} C. In our study sites, there is no freshwater flow or salinity gradient for most of the year; the creeks are small and open directly into the open ocean, with large tides and waves effectively mixing waters, meaning that at least for the dry season, the relationships between estimated mangrove cover and consumer $\delta^{13}C$ were only due to the presence of mangrove material and salinity had a limited effect.

If there was a measurable effect of mangrove-derived DIC- δ^{13} C on the time-averaged δ^{13} C of aquatic primary producers, this effect would be stronger in the upper reaches of the creeks and less in the lower reaches because close to the creek mouths, the water mixes more effectively. So, δ^{13} C of aquatic primary producers in the lower reaches would be more similar between sites, and if C₃ sources did not have any contribution to diets, no relationship between estimated mangrove cover and consumer $\delta^{13}C$ would be found for consumers collected at the lower reaches. This was however not the case (see Fig. 4). On the other hand, due to the lack of freshwater flow, more mangrove material would be accumulated in the creek beds during the dry season, meaning that there would be a higher availability of ¹³C-depleted mangrove carbon at this time, with a stronger effect on δ^{13} C of aquatic producers. If the detected differences in δ^{13} C were a result of differences in mangrove cover solely due to this indirect effect, then consumers should have lower $\delta^{13}C$ values in the dry season. While this was true for the banana prawn *P. merguiensis*, the opposite was true for the two fish species, including the planktivore A. vachellii, despite a previous study showing a positive relationship between DIC- δ^{13} C and planktivorous fish, which was not present for other trophic guilds (Abrantes et al. 2013).

Note also that MPB and (for Cocoa and Crab Creeks) seagrass have typically high δ^{13} C values, higher than plankton and generally more similar to C₄ grasses (Clementz and Koch 2001). So, any possible spatial differences in their δ^{13} C between estuaries would not lead to differences in results relating to the relative importance of C₃ sources, as these were well separated in δ^{13} C when compared to all other potential sources. Nevertheless, the high variability in source δ^{13} C used in the mixing model inputs (SD of ±3 ‰ for MPB and ±2 ‰ for the remaining sources),

coupled with the 1.5 ‰ uncertainty in TEF values, accounted for the uncertainty in source δ^{13} C resulting from the lack of local data on primary producer δ^{13} C and therefore the relative contribution of C₃ sources presented here can be considered conservative.

Differences in aquatic productivity between systems could have influenced the difference in importance of the different sources to consumers, but no productivity data were collected. Planktonic productivity is however likely to be similar between systems due to similarity in climate, environmental settings such as shading and depth and effective tidal mixing, while benthic productivity could differ between estuaries due to differences in area available for benthic production. However, the shallower estuaries, i.e., the estuaries with the largest intertidal area available for MPB production, were also those with denser and more extensive mangrove forests, so if MPB were of greater importance at these sites, then the contribution of C₃ sources would be relatively low, and this was not the case. Although the biomass of benthic algae in mangrove forests is generally low due to shading, these producers can be important in estuaries with greater areas of exposed habitat such as salt marshes, mudflats, and salt flats (Alongi 1988). In Australia's wet-dry tropics, these habitats are generally found at higher elevations and are less frequently inundated than mangroves, so for most of the time MPB are subjected to high temperatures, high salinities, and to desiccation, limiting productivity (Blanchard et al. 1996). Hence, differences in plankton and/or MPB productivity cannot explain the differences in C₃ contribution between estuaries.

It is also possible that other sources such as epiphytes growing on mangrove roots are important but are missing from the models. However, epiphytes are not likely to constitute important source for consumers in these estuaries as the close canopy of mangroves limits light penetration and, consequently, algal biomass and productivity. Also, the high tidal amplitude (maximum tidal range of ~4 m) means that for most of the time, these algae are subjected to desiccation or submerged in the waters generally turbid due to resuspension of soft sediment with the large tides, and both these factors limit the photosynthetic activity and productivity of epiphytic algae. Indeed, previous studies have shown that the abundance and productivity of algae in Australian wet and wet-dry tropical mangrove forests is low and that these areas are zones of net heterotrophy (Alongi et al. 1993; Alongi 1994). Nevertheless, epiphytes in these estuaries would likely have δ^{13} C close to plankton (e.g., Boon et al. 1997; Abrantes and Sheaves 2009a, 2009b; Nyunja et al. 2009; Al-Maslamani et al. 2013), and therefore, their inclusion in the models would not have affected the calculated contributions of C₃ sources and, therefore, the main conclusions of this study.

It could also be argued that the measured consumer $\delta^{13}C$ values are not a good representation of the average $\delta^{13}C$ values of the three species sampled, as these were based on only one analysed stable isotope sample per site per reach. However, each sample was composed of up to 15 individuals, and previous studies demonstrated that the analysis of $\sim 5-6$ individuals is sufficient to estimate mean δ^{13} C for estuarine prawns (Fry 1981) and fish (Mazumder et al. 2008) within an area. Furthermore, in the calculation of mean δ^{13} C, there is a complete agreement between mean $\delta^{13}C$ calculated using a number of individuals analysed separately and $\delta^{13}C$ calculated based on one sample composed by the same number of pooled individuals (Fry 1981). This means that the δ^{13} C value of one sample composed by 15 individuals combined is not different to the average δ^{13} C calculated based on 15 individuals analysed separately, and therefore, the measurement of individuals separately would not provide more information. Note also that the δ^{13} C variability of estuarine fish and invertebrates in North Queensland is generally low: of 67 fish and 4 penaeid species collected at various times from 35 systems in Central and North Queensland, $\delta^{13}C$ standard deviations ranged from 0.3 to 1.3 (25th–75th percentiles; n=273 for fish and n=56 for prawns) (authors' unpublished data). In those studies, the average number of replicates per species was only three, meaning that the SDs of δ^{13} C from up to 15 individuals is likely to be lower. Note also that Siarsolo was used in these models, as appropriate for models run based on one data point.

Seasonal Variability in Importance of C₃ Sources

Within each estuary, all three species ultimately relied on similar combinations of sources throughout the length of the estuaries in the wet season, but in the dry season, there were often large differences in the ultimate sources of nutrition used by different species and in different reaches. It is possible that this is related to the higher availability of nutrients in the wet season which resulted from the transport of material from upstream and the adjacent catchment with the freshwater flows, stimulating aquatic primary and secondary production (e.g., Hoover et al. 2006; Schlacher et al. 2008). More nutritive and easily assimilated material (i.e., plankton) would then be readily available, supporting abundant invertebrate communities that are prey for fish and other invertebrates. For example, in a study in Alligator Creek (located between Cocoa and Sandfly Creeks), a strong seasonality in density of zooplankton community was found, with much higher densities in the wet season than in the dry season (Robertson et al. 1988). Furthermore, different habitats and sites had relatively similar zooplankton communities in the wet season, but these differed in the dry season (Robertson et al. 1988). Since estuarine consumers can switch their diet to feed on temporarily abundant prey (Robertson et al. 1988; Baker and Sheaves 2009), the different species could feed on this abundant and similar prey assemblage at this time (Robertson et al. 1988), ending up with similar stable isotope composition. In the dry season, however, nutrient and food availability would be lower, and the assemblage of available prey would be less homogeneous throughout the length of the estuaries (Robertson et al. 1988), so the different species would have more diversified diets, feed on different prey assemblages at the different sites, and this would be reflected on differences in δ^{13} C between species and reaches. Further studies should be done to investigate this hypothesis.

The argument of increased productivity driven by nutrient input during the wet season may be seen as contradictory to the previously presented hypothesis of lack of significant effect of ¹³C-depleted DIC of mangrove origin over aquatic producers due to the effective flushing of these estuaries, i.e., should not this flux also flush out nutrients from the systems, particularly during the wet season when flows are higher? However, while DIC is likely more effectively flushed from these systems, a significant part of the heavier mangrove detritus probably settles and accumulates in the creek beds, where it becomes available to detritivores. The gentle topography of these creeks facilitates retention of this material. There are however no estimates of dissolved organic or inorganic carbon (DIC, DOC) or detritus residency times and exports for these small wet–dry tropical estuaries.

There was also evidence of seasonality in sources of nutrition for the three species. Interestingly, the different species had different patterns of seasonal change in importance of C₃ sources: while for A. vachellii and L. equulus, C3 sources were more important in the dry season, for P. merguiensis, C₃ sources were generally more important in the wet season. This could be because the different species are part of different food chains. For the two fish species, the lower importance of C₃ material during the wet season could have been a result of a more abundant small invertebrate prey community due to the increase in aquatic productivity that resulted from the input of nutrients with the wet season, as explained above. For example, although zooplankton assimilates both phytoplankton and detritus, it feeds selectively, preferring phytoplankton (Cole et al. 2006; Schlacher et al. 2009), so an increased phytoplankton productivity would lead to an increase in importance of aquatic sources and, consequently, in a decrease in relative importance of C₃ sources for these species and their predators. For P. merguiensis, the greater importance of C₃ material in the wet season could result from a greater input of mangrove detritus into the estuaries, as mangrove productivity and litterfall in this region is higher in the wet season (Robertson et al. 1988; Clough 1998). Unlike the two carnivorous fish species, P. merguiensis juveniles are mostly detritivorous (Robertson 1988), so higher availability of mangrove carbon would be more rapidly reflected into an increase in importance of mangrove carbon for the nutrition of this species. This explains the increase in importance of C₃ carbon for *P. merguiensis* in the lower and middle reaches, where most mangrove areas are concentrated. Although it is likely that increased mangrove productivity during the wet season is

somewhat offset by the reduced residence time due to higher flows, and that the detrital pool contains a higher proportion of algal matter at this time; results suggest that these effects are not sufficient to counteract the higher relative availability of mangrove detritus for detritivorous species during the wet season.

Therefore, it is possible that wet seasons have two different effects over these food webs, depending on the trophic ecology of the different species: the input of fresh nutrients stimulates aquatic productivity, fuelling algae-based food chains and reducing the relative importance of mangrove carbon for carnivores like *A. vachellii* and *L. equulus*, while the increase in available mangrove detritus due to increase mangrove productivity leads to an increase in importance of mangrove material for species that rely mostly on detritus-based food chains. This agrees with previous studies (mostly on freshwater systems) that show that detritivorous species are generally more affected by introduction of detrital material into a system than species that ultimately rely mostly on aquatic producers (Marczak et al. 2007; Abrantes and Sheaves 2010). However, further studies need to be conducted to test for this possibility.

The substantial importance of mangrove material detected in the present study is not in agreement with other studies in tropical regions, as most found limited importance of mangrove carbon to estuarine consumers (e.g., Fry and Ewel 2003; Layman 2007; Igulu et al. 2013 and references therein). Most studies found mangrove material to be important only for consumers within or in close proximity to the mangrove forests (e.g., Rodelli et al. 1984; Newell et al. 1995; Nyunja et al. 2009; Vaslet et al. 2012), especially in permanently inundated forests (Igulu et al. 2013). However, most of those studies were conducted in systems very different to those of the present study. For example, Heithaus et al. (2011) sampled an open coast area (Shark Bay, Western Australia) with low mangrove productivity (fringing mangroves) and with adjacent seagrass beds, so the potential for mangrove contribution was smaller. Indeed, most available studies were done in areas with adjacent productive seagrass beds (e.g., Loneragan et al. 1997; Nagelkerken and van der Velde 2004; Heithaus et al. 2011) and/or in much larger systems (e.g., Chanton and Lewis 2002; Abrantes et al. 2013) where mangrove detritus can be more easily diluted. Only a few recent studies have been conducted in areas where mangrove areas are not in close proximity to other productive coastal habitats such as seagrass beds, which can provide alternative food sources (Giarrizzo et al. 2011; Zagars et al. 2013). In those studies, like in the present study, mangrove carbon was found to be important for estuarine fish and invertebrate nutrition (Giarrizzo et al. 2011, Zagars et al. 2013).

Few studies have considered the seasonality in importance of terrestrial material transported from river catchments for tropical estuarine food webs. Those available suggest that this allochthonous source is seasonally important for aquatic consumers. For example, in bays and estuaries of Hong Kong, southern China (Wai et al. 2008, 2011), and Florida, USA (Chanton and Lewis 2002), in floodplain pools in North Queensland (Abrantes and Sheaves 2010), and in east African estuaries (Abrantes et al. 2013), there was a significant increase in importance of terrestrial material transported from the catchment during wet season. However, those studies considered systems very different to those from the present study: the North Queensland floodplain pools studied by Abrantes and Sheaves (2010) are small, relatively isolated and typically with a very narrow band of mangrove vegetation, so terrestrial organic matter transported from the catchment is likely to contribute a large proportion to the pool of available sources. The Hong Kong bays (Wai et al. 2008) receive large amounts of water from several hill streams that run through shrubland and forest during the wet summer monsoon, unlike the sites from the present study where rainfall is much lower, even in the wet season, and where the topography is much flatter. The Hong Kong (Wai et al. 2011), Florida (Chanton and Lewis 2002), and African estuaries (Abrantes et al. 2013), on the other hand, were much larger systems, with much larger catchments and discharges, so great quantities of terrestrial organic matter could be transported from their catchments, making a large contribution to aquatic food webs. Therefore, material from the catchment was likely to contribute to a much larger proportion of the total available carbon than for the systems considered in the present study, where small catchments and little rainfall during most of the year mean that there is limited potential for transport of terrestrial organic matter into the aquatic environment. Thus, unlike in perennial river systems with large catchments, estuarine food webs in small wet-dry tropical estuaries are likely to be less affected by impacts in the terrestrial environment landward of mangrove forests.

Conclusion

This study shows that mangroves are important contributors to estuarine food webs in small wet–dry tropical estuaries. In systems where extensive mangrove forests are present, mangrove-derived carbon can be the main source of nutrients supporting food webs. This is unlike in large perennial river systems, where aquatic sources such as plankton and benthic algae can have a greater importance (e.g., Chanton and Lewis 2002), most likely due to differences in ratio of mangrove to intertidal and open water area between these contrasting systems. There were also seasonal differences in sources of nutrition for food webs. Results suggest that this is, at least in part, due to the input of nutrients during the wet season, which stimulated algae-based food chains, reducing the relative importance of mangrove carbon for carnivorous fish like *A. vachellii* and *L. equulus*. At the same time, increases in

mangrove productivity during the hot wet season seemed to lead to increases in importance of mangrove material for detritus-based food chains.

Acknowledgments We thank the many volunteers, in particular A. Johnson for field assistance. This research was supported by an Australian Government Marine and Tropical Sciences Research Facility (MTSRF) grant to MS and RMC, and by a Winifred Violet Scott Foundation grant to KGA. Work was conducted in accordance with institutional, national, and international guidelines concerning the use of animals in research, under the Ethics Permit A1210 from James Cook University. We also thank the anonymous reviewers for their insightful comments.

References

- Abrantes, K., and M. Sheaves. 2008. Incorporation of terrestrial wetland material into aquatic food webs in a tropical estuarine wetland. *Estuarine, Coastal and Shelf Science* 80: 401–412.
- Abrantes, K., and M. Sheaves. 2009a. Food web structure in a nearpristine mangrove area of the Australian Wet Tropics. *Estuarine, Coastal and Shelf Science* 82: 597–607.
- Abrantes, K., and M. Sheaves. 2009b. Sources of nutrition supporting juvenile penaeid prawns in an Australian dry tropics estuary. *Marine* and Freshwater Research 60: 949–959.
- Abrantes, K.G., and M. Sheaves. 2010. Importance of freshwater flow in terrestrial-aquatic energetic connectivity in intermittently connected estuaries of tropical Australia. *Marine Biology* 157: 2071–2086.
- Abrantes, K. G, Barnett, A, Marwick, T. R, Bouillon S. 2013. Importance of terrestrial subsidies for estuarine food webs in contrasting east African catchments. Ecosphere 4:Art14
- Al-Maslamani, I., M. Walton, H. Kennedy, M. Al-Mohannadi, and L. Le Vay. 2013. Are mangroves in arid environments isolated systems? Life-history and evidence of dietary contribution from inwelling in a mangrove-resident shrimp species. *Estuarine, Coastal and Shelf Science* 124: 56–63.
- Alongi, D.M. 1988. Bacterial productivity and microbial biomass in tropical mangrove sediments. *Microbial Ecology* 15: 59–79.
- Alongi, D.M. 1994. Zonation and seasonality of benthic primary production and community respiration in tropical mangrove forests. *Oecologia* 98: 320–327.
- Alongi, D.M., P. Christoffersen, and F. Tirendi. 1993. The influence of forest type on microbial-nutrient relationships in tropical mangrove sediments. *Journal of Experimental Marine Biology and Ecology* 171: 201–223.
- Baker, R., and M. Sheaves. 2009. Refugees or ravenous predators: detecting predation on new recruits to tropical estuarine nurseries. *Wetlands Ecology and Management* 17: 317–330.
- Barnes, C., C.J. Sweeting, S. Jennings, J.T. Barry, and N.V.C. Polunin. 2007. Effect of temperature and ration size on carbon and nitrogen stable isotope trophic fractionation. *Functional Ecology* 21: 356– 362.
- Beck, M.W., K.L. Heck Jr., K.W. Able, D.L. Childers, D.B. Eggleston, B.M. Gillanders, B. Halpern, C.G. Hays, K. Hoshino, T.J. Minello, R.J. Orth, P.F. Sheridan, and M.P. Weinstein. 2001. The identification, conservation, and management of estuarine and marine nurseries for fish and invertebrates. *BioScience* 51: 633–641.
- Blanchard, G.F., J.-M. Guarini, P. Richard, P. Gros, and F. Mornet. 1996. Quantifying the short-term temperature effect on light-saturated photosynthesis of intertidal microphytobenthos. *Marine Ecology Progress Series* 134: 309–313.

- Bond, A.L., and A.W. Diamond. 2011. Recent Bayesian stable-isotope mixing models are highly sensitive to variation in discrimination factors. *Ecological Applications* 21: 1017–1023.
- Boon, P.I., F.L. Bird, and S.E. Bunn. 1997. Diet of the intertidal callianassid shrimps *Biffarius arenosus* and *Trypea australiensis* (Decapoda:Thalassinidea) in Western Port (southern Australia), determined with multiple stable-isotope analyses. *Marine and Freshwater Research* 48: 503–511.
- Bouillon, S., P.C. Mohan, N. Sreenivas, and F. Dehairs. 2000. Sources of suspended organic matter and selective feeding by zooplankton in an estuarine mangrove ecosystem as traced by stable isotopes. *Marine Ecology Progress Series* 208: 79–92.
- Bouillon, S., T. Moens, I. Overmeer, N. Koedam, and F. Dehairs. 2004. Resource utilization patterns of epifauna from mangrove forests with contrasting inputs of local versus imported organic matter. *Marine Ecology Progress Series* 278: 77–88.
- Bouillon, S., J.J. Middelburg, F. Dehairs, A.V. Borges, G. Abril, M.R. Flindt, S. Ulomi, and E. Kristensen. 2007. Importance of intertidal sediment processes and porewater exchange on the water column biogeochemistry in a pristine mangrove creek (Ras Dege, Tanzania). *Biogeosciences* 4: 311–322.
- Bouillon, S., R.M. Connolly, and S.Y. Lee. 2008. Organic matter exchange and cycling in mangrove ecosystems: recent insights from stable isotope studies. *Journal of Sea Research* 59: 44–58.
- Bouillon, S., R.M. Connolly, and D.P. Gillikin. 2011. Use of stable isotopes to understand food webs and ecosystem functioning in estuaries. In *Treatise on Estuarine and Coastal Science*, ed. E. Wolanski and D.S. McLusky. Waltham: Academic Press.
- Boys, C.A., F.J. Kroon, T.M. Glasby, and K. Wilkinson. 2012. Improved fish and crustacean passage in tidal creeks following floodgate remediation. *Journal of Applied Ecology* 49: 223–233.
- Buchheister, A., and R.J. Latour. 2010. Turnover and fractionation of carbon and nitrogen stable isotopes in tissues of a migratory coastal predator, summer flounder (*Paralichthys dentatus*). Canadian Journal of Fisheries and Aquatic Sciences 67: 445–461.
- Caut, S., E. Angulo, and F. Courchamp. 2009. Variation in discrimination factors (Δ^{15} N and Δ^{13} C): the effect of diet isotopic values and applications for diet reconstruction. *Journal of Animal Ecology* 46: 443–453.
- Cerling, T.E., J.M. Harris, B.J. MacFadden, M.G. Leakey, J. Quade, V. Eisenmann, and J.R. Ehleringer. 1997. Global vegetation change through the Miocene/Pliocene boundary. *Nature* 389: 153–158.
- Chanton, J., and F.G. Lewis. 2002. Examination of coupling between primary and secondary production in a river-dominated estuary: Apalachicola Bay, Florida, USA. *Limnology and Oceanography* 47: 683–697.
- Chong, V.C., C.B. Low, and T. Ichikawa. 2001. Contribution of mangrove detritus to juvenile prawn nutrition: a dual stable isotope study in a Malaysian mangrove forest. *Marine Biology* 138: 77–86.
- Clementz, M.T., and P.L. Koch. 2001. Differentiating aquatic mammal habitat and foraging ecology with stable isotopes in tooth enamel. *Oecologia* 129: 461–472.
- Cloern, J.E., E.A. Canuel, and D. Harris. 2002. Stable carbon and nitrogen isotope composition of aquatic and terrestrial plants of the San Francisco Bay estuarine system. *Limnology and Oceanography* 47: 713–729.
- Clough, B. 1998. Mangrove forest productivity and biomass accumulation in Hinchinbrook Channel, Australia. *Mangroves and Salt Marshes* 2: 191–198.
- Cole, J.J., S.R. Carpenter, M.L. Pace, M.C. Van de Bogert, J.L. Kitchell, and J.R. Hodgson. 2006. Differential support of lake food webs by three types of terrestrial organic carbon. *Ecology Letters* 9: 558–568.
- Dahlgren, C.P., G. Kellison, A.J. Adams, B.M. Gillanders, M.S. Kendall, C.A. Layman, J.A. Ley, I. Nagelkerken, and J.E. Serafy. 2006. Marine nurseries and effective juvenile habitats: concepts and applications. *Marine Ecology Progress Series* 312: 291–295.

- De'ath, G., and K.E. Fabricius. 2000. Classification and regression trees: a powerful yet simple technique for ecological data analysis. *Ecology* 81: 3178–3192.
- DeNiro, M.J., and S. Epstein. 1978. Influence of diet in the distribution of carbon isotopes in animals. *Geochimica et Cosmochimica Acta* 42: 495–506.
- Elsdon, T.S., S. Ayvazian, K.W. McMahon, and S.R. Thorrold. 2010. Experimental evaluation of stable isotope fractionation in fish muscle and otoliths. *Marine Ecology Progress Series* 408: 195–205.
- Erskine, W.D., M.J. Saynor, L. Erskine, K.G. Evans, and D.R. Moliere. 2005. A preliminary typology of Australian tropical rivers and implications for fish community ecology. *Marine and Freshwater Research* 56: 253–267.
- Finlayson, B., and T. McMahon. 1988. Australia vs the world: a comparative analysis of streamflow characteristics. In *Fluvial Geomorphology* of Australia, ed. R. Werner. Sydney: Academic Press.
- France, R.L. 1996. Scope for use of stable carbon isotopes in discerning the incorporation of forest detritus into aquatic foodwebs. *Hydrobiologia* 325: 219–222.
- Fry, B. 1981. Natural stable carbon isotope tag traces Texas shrimp migrations. *Fishery Bulletin* 79: 337–345.
- Fry, B. 2002. Conservative mixing of stable isotopes across estuarine salinity gradients: a conceptual framework for monitoring watershed influences on downstream fisheries production. *Estuaries* 25: 264– 271.
- Fry, B., and K.C. Ewel. 2003. Using stable isotopes in mangrove fisheries research—a review and outlook. *Isotopes in Environmental and Health Studies* 39: 191–196.
- Giarrizzo, T., R. Schwamborn, and U. Saint-Paul. 2011. Utilization of carbon sources in a northern Brazilian mangrove ecosystem. *Estuarine Coastal and Shelf Science* 95: 447–457.
- Gillikin, D.P., A. Lorrain, S. Bouillon, P. Willenz, and F. Dehairs. 2006. Stable carbon isotopic composition of *Mytilus edulis* shells: relation to metabolism, salinity, δ¹³C-DIC and phytoplankton. *Organic Geochemistry* 37: 1371–1382.
- Gorokhova, E., and S. Hansson. 1999. An experimental study on variations in stable carbon and nitrogen fractionation during growth of *Mysis mixta* and *Neomysis integer. Canadian Journal of Fisheries* and Aquatic Sciences 56: 2203–2210.
- Guelinckx, J., J. Maes, P. van den Driessche, B. Geysen, F. Dehairs, and F. Ollevier. 2007. Changes in δ¹³C and δ¹⁵N in different tissues of juvenile sand goby *Pomatoschistus minutus*: a laboratory dietswitch experiment. *Marine Ecology Progress Series* 341: 205–215.
- Guest, M., R.M. Connolly, and N. Loneragan. 2004. Within and amongsite variability in δ^{13} C and δ^{15} N for three estuarine producers, *Sporobolus virginicus, Zostera capricorni*, and epiphytes of *Z. capricorni. Aquatic Botany* 79: 87–94.
- Haywood, M.D.E., and D.J. Staples. 1993. Field estimates of growth and mortality of juvenile banana prawns (*Penaeus merguiensis*). *Marine Biology* 116: 407–416.
- Heithaus, E.R., P.A. Heithaus, M.R. Heithaus, D. Burkholder, and C.A. Layman. 2011. Trophic dynamics in a relatively pristine subtropical fringing mangrove community. *Marine Ecology Progress Series* 428: 49–61.
- Hemminga, M.A., and M.A. Mateo. 1996. Stable carbon isotopes in seagrasses: variability in ratios and use in ecological studies. *Marine Ecology Progress Series* 140: 285–298.
- Hoover, R.S., D. Hoover, M. Miller, M.R. Landry, E.H. DeCarlo, and F.T. Mackenzie. 2006. Zooplankton response to storm runoff in a tropical estuary: bottom-up and top-down controls. *Marine Ecology Progress Series* 318: 187–201.
- Igulu, M., I. Nagelkerken, G. van der Velde, and Y. Mgaya. 2013. Mangrove fish production is largely fuelled by external food sources: a stable isotope analysis of fishes at the individual, species, and community levels from across the globe. *Ecosystems* 16: 1336–1352.

- Jennerjahn, T.C., and V. Ittekkot. 2002. Relevance of mangroves for the production and deposition of organic matter along tropical continental margins. *Naturwissenschaften* 89: 23–30.
- Kristensen, E., S. Bouillon, T. Dittmar, and C. Marchand. 2008. Organic carbon dynamics in mangrove ecosystems: a review. *Aquatic Botany* 89: 201–219.
- Lancaster, J., and S. Waldron. 2001. Stable isotope values of lotic invertebrates: sources of variation, experimental design, and statistical interpretation. *Limnology and Oceanography* 46: 723–730.
- Layman, C.A. 2007. What can stable isotope ratio reveal about mangroves as fish habitat? *Bulletin of Marine Science* 80: 513–527.
- Lin, H.-J., W.-Y. Kao, and Y.-T. Wang. 2007. Analyses of stomach contents and stable isotopes reveal food sources of estuarine detritivorous fish in tropical/subtropical Taiwan. *Estuarine, Coastal and Shelf Science* 73: 527–537.
- Loneragan, N.R., S.E. Bunn, and D.M. Kellaway. 1997. Are mangroves and seagrasses sources of organic carbon for penaeid prawns in a tropical Australian estuary? A multiple stable-isotope study. *Marine Biology* 130: 289–300.
- Lotze, H.K., H.S. Lenihan, B.J. Bourque, R.H. Bradbury, R.G. Cooke, M.C. Kay, S.M. Kidwell, M.X. Kirby, C.H. Peterson, and J.B.C. Jackson. 2006. Depletion, degradation, and recovery potential of estuaries and coastal seas. *Science* 312: 1806–1809.
- Maher, D.T., I.R. Santos, L. Golsby-Smith, J. Gleeson, and B.D. Eyre. 2013. Groundwater-derived dissolved inorganic and organic carbon exports from a mangrove tidal creek: the missing mangrove carbon sink? *Limnology and Oceanography* 58: 475–488.
- Marczak, L.B., R.M. Thompson, and J.S. Richardson. 2007. Metaanalysis: trophic level, habitat, and productivity shape the food web effects of resource subsidies. *Ecology* 88: 140–148.
- Mavuti, K.M., J.A. Nyunja, and E.O. Wakwabi. 2007. Trophic ecology of some common juvenile fish species in Mtwapa Creek, Kenya. West Indian Ocean Journal of Marine Science 3: 179–188.
- Mazumder, D., R.J. Williams, D. Reir, N. Saintilan, and R. Szymczak. 2008. Variability of stable isotope ratios of glassfish (*Ambassis jacksoniensis*) from mangrove/saltmarsh environments in southeast Australia and emplications for choosing sample size. *Environmental Bioindicators* 3: 114–123.
- McCutchan, J.H., W.M. Lewis Jr., C. Kendall, and C.C. McGrath. 2003. Variation in trophic shift for stable isotope ratios of carbon, nitrogen and sulfur. *Oikos* 102: 378–390.
- Nagelkerken, I., and G. van der Velde. 2004. Are Caribbean mangroves important feeding grounds for juvenile reef fish from adjacent seagrass beds? *Marine Ecology Progress Series* 274: 143–151.
- Newell, R.I.E., N. Marshall, A. Sasekumar, and V.C. Chong. 1995. Relative importance of benthic microalgae, phytoplankton, and mangroves as sources of nutrition for penaeid prawns and other coastal invertebrates from Malaysia. *Marine Biology* 123: 595– 606.
- Nyunja, J., M. Ntiba, J. Onyari, K. Mavuti, K. Soetaert, and S. Bouillon. 2009. Carbon sources supporting a diverse fish community in a tropical coastal ecosystem (Gazi Bay, Kenya). *Estuarine, Coastal* and Shelf Science 83: 333–341.
- Parnell, A.C., R. Inger, S. Bearhop, and A.L. Jackson. 2010. Source partitioning using stable isotopes: coping with too much variation. *PLoS ONE* 5: e9672.
- Peel, M.C., B.L. Finlayson, and T.A. McMahon. 2007. Updated world map of the Köppen-Geiger climate classification. *Hydrology and Earth System Sciences Discussions* 4: 439–473.
- Peterson, B.J., R.W. Howarth, and R.H. Garritt. 1986. Sulfur and carbon isotopes as tracers of salt-marsh organic matter flow. *Ecology* 67: 865–874.
- Polis, G.A., W.B. Anderson, and R.D. Holt. 1997. Towards an integration of landscape and food web ecology: the dynamics of spatially subsidized food webs. *Annual Review of Ecology and Systematics* 28: 289–316.

- Post, D.M. 2002. Using stable isotopes to estimate trophic position: models, methods and assumptions. *Ecology* 83: 703–718.
- Post, D.M., C.A. Layman, D.A. Arrington, G. Takimoto, J. Quattrochi, and C.G. Montanã. 2007. Getting to the fat of the matter: models, methods and assumptions for dealing with lipids in stable isotope analyses. *Oecologia* 152: 179–189.
- Ridd, P.V., and T. Stieglitz. 2002. Dry season salinity changes in arid estuaries fringed by mangroves and saltflats. *Estuarine, Coastal and Shelf Science* 54: 1039–1049.
- Ridd, P., R. Sam, S. Hollins, and G. Brunskill. 1997. Water, salt and nutrient fluxes of tropical tidal salt flats. *Mangroves and Salt Marshes* 1: 229–238.
- Robertson, A.I. 1988. Abundance, diet and predators of juvenile banana prawns, *Penaeus merguiensis*, in a tropical mangrove estuary. *Australian Journal of Marine & Freshwater Research* 39: 467–478.
- Robertson, A.I., and S.J.M. Blaber. 1992. Plankton, epibenthos and fish communities. In *Tropical Mangrove Ecossystems, Book 41*, ed. A.I. Robertson and D.M. Alongi. Washington, DC: American Geophysical Union.
- Robertson, A.I., and N.C. Duke. 1990. Recruitment, growth and residence time of fishes in a tropical Australian mangrove system. *Estuarine, Coastal and Shelf Science* 31: 723–743.
- Robertson, A.I., P. Dixon, and P.A. Daniel. 1988. Zooplankton dynamics in mangrove and other nearshore habitats in tropical Australia. *Marine Ecology Progress Series* 43: 139–150.
- Rodelli, M.R., J.N. Gearing, P.J. Gearing, N. Marshall, and A. Sasekumar. 1984. Stable isotope ratio as a tracer of mangrove carbon in Malaysian ecosystems. *Oecologia* 61: 326–333.
- Schlacher, T.A., A.J. Skillington, R.M. Connolly, W. Robinson, and T.F. Gaston. 2008. Coupling between marine plankton and freshwater flow in the plumes off a small estuary. *International Review of Hydrobiology* 93: 641–658.
- Schlacher, T.A., R.M. Connolly, A.J. Skillington, and T.F. Gaston. 2009. Can export of organic matter from estuaries support zooplankton in nearshore, marine plumes? *Aquatic Ecology* 43: 383–393.
- Sheaves, M. 1996. Do spatial differences in the abundance of two serranid fishes in estuaries of tropical australia reflect long-term salinity patterns? *Marine Ecology Progress Series* 137: 39–49.
- Sheaves, M., and R. Johnston. 2009. Ecological drivers of spatial variability among fish fauna of 21 tropical Australian estuaries. *Marine Ecology Progress Series* 385: 245–260.
- Sheaves, M., and R. Johnston. 2010. Implications of spatial variability of fish assemblages for monitoring of Australia's tropical estuaries. *Aquatic Conservation: Marine and Freshwater Ecosystems* 20: 348–356.
- Sheaves, M., and B. Molony. 2000. Short-circuit in the mangrove food chain. *Marine Ecology Progress Series* 199: 97–109.
- Sheaves, M., R. Johnston, and R. Connolly. 2010. Temporal dynamics of fish assemblages of natural and artificial tropical estuaries. *Marine Ecology Progress Series* 410: 143–157.

- Sheaves, M., R. Johnston, and R.M. Connolly. 2012. Fish assemblages as indicators of estuary ecosystem health. *Wetlands Ecology and Management* 20: 477–490.
- Sheaves, M., R. Johnston, A. Johnson, R. Baker, and R. Connolly. 2013. Nursery function drives temporal patterns in fish assemblage structure in four tropical estuaries. *Estuaries and Coasts* 36: 893–905.
- Sheaves, M., J. Brookes, R. Coles, M. Freckelton, P. Groves, R. Johnston, and P. Winberg. 2014. Repair and revitalisation of Australia's tropical estuaries and coastal wetlands: opportunities and constraints for the reinstatement of lost function and productivity. *Marine Policy* 47: 23–38.
- Suring, E., and S.R. Wing. 2009. Isotopic turnover rate and fractionation in multiple tissues of red rock lobster (*Jasus edwardsii*) and blue cod (*Parapercis colias*): consequences for ecological studies. *Journal of Experimental Marine Biology and Ecology* 370: 56–63.
- Vance, D.J., M.D.E. Haywood, D.S. Heales, R.A. Kenyon, N.R. Loneragan, and R.C. Pendrey. 1996. How far do prawns and fish move into mangroves? Distribution of juvenile banana prawns *Penaeus merguiensis* and fish in a tropical mangrove forest in northern Australia. *Marine Ecology Progress Series* 131: 115–124.
- Vander Zanden, M.J., and J.B. Rasmussen. 2001. Variation in δ^{15} N and δ^{13} C trophic fractionation: implication for aquatic food web studies. *Limnology and Oceanography* 46: 2061–2066.
- Vaslet, A., D.L. Phillips, C. France, I.C. Feller, and C.C. Baldwin. 2012. The relative importance of mangroves and seagrass beds as feeding areas for resident and transient fishes among different mangrove habitats in Florida and Belize: evidence from dietary and stableisotope analyses. *Journal of Experimental Marine Biology and Ecology* 434–435: 81–93.
- Wai, T.-C., J.S. Ng, K.M. Leung, D. Dudgeon, and G.A. Williams. 2008. The source and fate of organic matter and the significance of detrital pathways in a tropical coastal ecosystem. *Limnology and Oceanography* 53: 1479–1492.
- Wai, T.-C., K.M.Y. Leung, S.Y.T. Sin, A. Cornish, D. Dudgeon, and G.A. Williamsa. 2011. Spatial, seasonal, and ontogenetic variations in the significance of detrital pathways and terrestrial carbon for a benthic shark, *Chiloscyllium plagiosum* (Hemiscylliidae), in a tropical estuary. *Limnology and Oceanography* 56: 1035–1053.
- Weidel, B.C., S.R. Carpenter, J.F. Kitchell, and M.J. Vander Zanden. 2011. Rates and components of carbon turnover in fish muscle: insights from bioenergetics models and a whole-lake ¹³C addition. *Canadian Journal of Fisheries and Aquatic Sciences* 68: 387–399.
- Wilson, J.P., and M. Sheaves. 2001. Short-term temporal variations in taxonomic composition and trophic structure of a tropical estuarine fish assemblage. *Marine Biology* 139: 787–796.
- Yokoyama, H., A. Tamaki, K. Harada, K. Shimoda, K. Koyama, and Y. Ishihi. 2005. Variability of diet-tissue isotopic fractionation in estuarine macrobenthos. *Marine Ecology Progress Series* 296: 115–128.
- Zagars, M., K. Ikejima, A. Kasai, N. Arai, and P. Tongnunui. 2013. Trophic characteristics of a mangrove fish community in Southwest Thailand: Important mangrove contribution and intraspecies feeding variability. *Estuarine Coastal and Shelf Science* 119: 145–152.